# **R** documentation

# of all in 'QSutils/man'

# September 4, 2018

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#### **Description**

\packageDescriptionQSutils

#### **Details**

The DESCRIPTION file: \packageDESCRIPTIONQSutils

### Author(s)

\packageAuthorQSutils

Maintainer: \packageMaintainerQSutils

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83.doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

Collapse 3

Collapse reads into haplotypes and frequencies
--

### **Description**

Collapse summarizes aligned reads into haplotypes with their frequencies. Recollapse is used to update the collapse after some type of manipulation may have resulted in duplicate haplotypes.

### Usage

```
Collapse(seqs)
Recollapse(seqs,nr)
```

### Arguments

seqs DNAStringSet or AAStringSet object with the sequences to collapse.

nr Vector with the haplotype counts.

#### **Details**

Recollapse is used when haplotypes may become equivalent after some type of manipulation. It removes duplicate sequences and updates their frequencies.

#### Value

Collapse and Recollapse return a list with two elements.

nr Vector of the haplotype counts

hseqs DNAStringSet or AAStringSet with the haplotype sequence

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

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### **Examples**

ConsSeq

Consensus sequence given an alignment and frequencies

# Description

ConsSeq determines the consensus sequence from a set of haplotypes.

# Usage

```
ConsSeq(seqs, w=NULL)
```

# **Arguments**

seqs DNAStringSet or AAStringSet object with the haplotype sequences.

w An optional numeric vector with the haplotype counts.

#### **Details**

The most frequent nucleotide or amino acid at each position is taken. No IUPAC ambiguity codes are considered; in the case of ties, the consensus nucleotide is decided randomly.

# Value

Character vector with the consensus sequence.

### Author(s)

Josep Gregori and Mercedes Guerrero

### See Also

ReadAmplSeqs

CorrectGapsAndNs 5

### **Examples**

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
ConsSeq(lst$hseqs,lst$nr)</pre>
```

CorrectGapsAndNs

Function to correct an alignment with gaps and Ns

### **Description**

Corrects positions in a DNAStringSet or AAStringSet of aligned haplotypes, replacing gaps and Ns (indeterminates) with the nucleotide or amino acid from the corresponding position in the reference sequence.

### Usage

CorrectGapsAndNs(hseqs, ref.seq)

### **Arguments**

hseqs DNAStringSet or AAStringSet object with the alignment to correct.

ref.seq Character vector with the reference sequence of the alignment.

#### Value

DNAStringSet or AAStringSet object with the sequences corrected. Duplicate haplotypes may arise as a consequence of this operation. See Recollapse.

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

#### See Also

Recollapse

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### **Examples**

```
# Create a random reference sequence.
ref.seq <-GetRandomSeq(50)
ref.seq

# Create an alignment with gaps and Ns.
symb <- c(".","-","N")
nseqs <- 12
p <- c(0.9,0.06,0.04)
hseqs <- matrix(sample(symb,50*nseqs,replace=TRUE,prob=p),ncol=50)
hseqs <- apply(hseqs,1,paste,collapse="")
hseqs
hseqs <- DNAStringSet(hseqs)

# Apply the function and visualize the result.
cseqs <- CorrectGapsAndNs(hseqs,as.character(ref.seq))
c(ref.seq,as.character(cseqs))</pre>
```

DBrule

Genotyping by the DB rule

# **Description**

Computes the nearest cluster to a given sequence.

### Usage

```
DBrule(grpDist, hr, oDist, g.names = NULL)
```

### **Arguments**

grpDist Distances between reference sequences.

hr Factor or a vector of integers that contains the type or subtype for each reference

sequence.

oDist Distance from the sequence to be classified to the reference sequences.

g.names Type or subtype names to classify the sequence.

### Value

List with three elements:

Phi 2 Vector with the distances to each cluster.

DB.rule The index of the nearest cluster.

Type Name of the nearest cluster.

### Author(s)

Josep Gregori and Mercedes Guerrero

Diverge 7

### References

Caballero A, Gregori J, Homs M, Tabernero D, Gonzalez C, Quer J, Blasi M, Casillas R, Nieto L, Riveiro-Barciela M, Esteban R, Buti M, Rodriguez-Frias F. Complex Genotype Mixtures Analyzed by Deep Sequencing in Two Different Regions of Hepatitis B Virus. PLoS One. 2015 Dec 29;10(12):e0144816. doi: 10.1371/journal.pone.0144816. eCollection 2015. PubMed PMID: 26714168; PubMed Central PMCID: PMC4695080.

### **Examples**

```
# Load haplotype to be genotyped.
lst <- ReadAmplSeqs("QSutils/extdata/Unknown-Genotype.fna")</pre>
hseq <- lst$hseq[1]</pre>
# Load genotype references.
RefSeqs <- readDNAStringSet("QSutils/extdata/GenotypeStandards_A-H.fas")</pre>
# Compute pairwise distances.
dm <- as.matrix(DNA.dist(c(hseq,RefSeqs),model="K80"))</pre>
# Distances between genotype RefSeqs
dgrp <- dm[-1,-1]
grp <- factor(substr(rownames(dgrp),1,1))</pre>
hr <- as.integer(grp)</pre>
# Distance of the query haplotype to the reference sequence.
d <- dm[1,-1]
# Genotyping by the DB rule.
dsc <- DBrule(dgrp,hr,d,levels(grp))</pre>
dsc
```

Diverge

Generate a set of diverging haplotypes

# **Description**

Generates a set of diverging haplotypes from the given DNA sequence. The haplotypes produced share a pattern of divergence with an increasing number of mutations.

### Usage

```
Diverge(vm, seq)
```

# **Arguments**

vm Vector with number of diverging mutations to be generated. seq Reference sequence from which to generate the variants.

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### **Details**

max(vm) Positions in the given sequence are randomly generated. A substitution is also randomly produced for each of these positions. A haplotype is generated for each element in vm, so that it contains vm[i] substitutions of those previously generated.

#### Value

Character string vector with the segregating haplotypes generated.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### See Also

GetRandomSeg

### **Examples**

```
set.seed(123)
m1 <- GetRandomSeq(50)
hpl <- Diverge(3:6,m1)
DottedAlignment(DNAStringSet(hpl))</pre>
```

DNA.dist

Matrix of DNA distances given an alignment

# **Description**

Function to compute a matrix of pairwise distances from DNA sequences using a model of DNA evolution. It relies on the dist.dna() function in the APE package.

### Usage

```
DNA.dist(seqs, model = "raw", gamma = FALSE, pairwise.deletion = FALSE)
```

### **Arguments**

seqs DNAStringSet object with the aligned haplotypes.

model Evolutionary model to compute genetic distance by default "raw", but "N",

"TS", "TV", "JC69", "K80", "F81", "K81", "F84", "BH87", "T92","TN93",

"GG95", "logdet", "paralin", "indel", or "indelblock" can also be used.

gamma Gamma parameter possibly used to apply a correction to the distances or FALSE

(by default).

pairwise.deletion

A logical indicating whether to delete sites with missing data (gaps) in a pairwise manner. The default is to delete sites with at least one missing datum in all sequences.

DottedAlignment 9

#### Value

Object of class "dist" with pairwise distances.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Paradis E., Claude J. and Strimmer K., APE: analyses of phylogenetics and evolution in R language. Bioinformatics. 2004, 20, 289-290

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

```
dist.dna
```

### **Examples**

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
dst <- DNA.dist(lst$hseqs,model="N")
dst</pre>
```

DottedAlignment

Align haplotypes into a dotted alignment

### **Description**

Given an alignment, it takes the first sequence as reference, and depicts all equivalences in the alignment as dots, leaving only the variants with respect to the reference.

### Usage

```
DottedAlignment(hseqs)
```

### **Arguments**

hseqs

DNAStringSet or AAStringSet with haplotype sequences..

### Value

A character string vector of the alignment, with dots in the conserved positions.

DSFT

### Author(s)

Josep Gregori and Mercedes Guerrero

# See Also

ReadAmplSeqs

# **Examples**

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
strs <- DottedAlignment(lst$hseqs)

# Create a data frame to visualize the result.
data.frame(Hpl=strs,stringsAsFactors=FALSE)</pre>
```

**DSFT** 

Downsampling followed by fringe trimming

### **Description**

Diversity indices are influenced to a greater or lesser degree by the sample size on which they are computed. This function helps to minimize the bias inherent to sample size. First the vector of abundances is scaled to a smaller sample size, then all haplotypes with abundances below a given threshold are excluded with high confidence.

# Usage

```
DSFT(nr, size, p.cut = 0.002, conf = 0.95)
```

# **Arguments**

nr Vector of observed haplotype counts.

size Size to downsample.

p.cut Abundance threshold.

conf Confidence in trimming.

# Value

Vector of logicals, with false the haplotypes to be excluded.

#### Author(s)

Josep Gregori and Mercedes Guerrero

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### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

# **Examples**

```
# Generate viral quasispecies abundance data.
set.seed(123)
n <- 2000
y \leftarrow geom.series(n, 0.8) + geom.series(n, 0.0004)
nr.pop <- round(y*1e7)</pre>
# Get a sample of 10000 reads from this population.
sz2 <- 10000
nr.sz2 <- table(sample(length(nr.pop),size=sz2,replace=TRUE,p=nr.pop))</pre>
# Filter out haplotypes below 0.1%.
thr <- 0.1
f1 <- nr.sz2 >= sz2 * thr/100
nr.sz2 \leftarrow nr.sz2[f1]
Shannon(nr.sz2) #0.630521
# DSFT to 5000 reads.
sz1 <- 5000
fl <- DSFT(nr.sz2,sz1)</pre>
nr.sz2 \leftarrow nr.sz2[f1]
# Compute size corrected Shannon entropy.
Shannon(nr.sz2) #0.6189798
```

FAD

Functional attribute diversity

# Description

Computes the Functional Attribute Diversity as the sum of elements in the pairwise distance matrix.

### Usage

FAD(dst)

### **Arguments**

dst

A "dist" object or a symmetrical matrix with pairwise distances.

### Value

A value that corresponds to the Functional Attribute Diversity. The sum of matrix elements.

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### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

### See Also

DNA.dist

# **Examples**

```
# Create the object.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Compute the DNA distance matrix.
dst <- DNA.dist(lst$hseqs,model="N")
FAD(dst)</pre>
```

fn.ab.1

Vector of abundances by sequential powers of a fraction

### **Description**

Function to simulate haplotype abundances in the quasispecies by sequential powers of a fraction.

# Usage

```
fn.ab.1(n, h = 10000, r = 0.5)
```

# **Arguments**

n Number of counts to compute.

h Highest abundance value.

r A number in the range 0 < r < 1.

# **Details**

The abundances, as counts, are computed according to the following equation, taking the integer part:

$$max(hr^{(i-1)}, 1); 0 < r < 1; i = 1..n$$

The lower r, the faster the decrease in abundance.

fn.ab.2

# Value

Numeric vector with n decreasing counts, where the first element equals h, and no element is lower than 1

# Author(s)

Josep Gregori and Mercedes Guerrero

### See Also

```
fn.ab.2, fn.ab.3, geom.series ,GetRandomSeq, GenerateVars, Diverge
```

# **Examples**

```
# Simulate a quasispecies alignment.
m1 <- GetRandomSeq(50)
v1 <- GenerateVars(m1,50,2,c(10,1))
qs <- c(m1,v1)
w <- fn.ab.1(length(qs),h=1000,r=1.5)</pre>
```

fn.ab.2

Vector of abundances by a power of decreasing fractions

# Description

Function to simulate haplotype abundances in the quasispecies by a power of decreasing fractions.

# Usage

```
fn.ab.2(n, h = 10000, r = 3)
```

#### **Arguments**

n Number of counts to compute.

h Highest abundance value.

The power of the function. A value larger than 0, usually in the range 0.5 < r < 4.

#### **Details**

The abundances, as counts, are computed according to the following equation, taking the integer part:

$$\max(h\ \left(\frac{1}{i}\right)^r,1);\ r>0;\ i=1..n$$

The higher r, the faster the decrease in abundances.

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### Value

Numeric vector with n decreasing counts, where the first element equals h and no element is lower than 1

# Author(s)

Josep Gregori and Mercedes Guerrero

#### See Also

```
fn.ab.1, fn.ab.3, geom.series, GetRandomSeq, GenerateVars, Diverge
```

# **Examples**

```
# Simulate a quasispecies alignment.
m1 <- GetRandomSeq(50)
v1 <- GenerateVars(m1,50,2,c(10,1))
qs <- c(m1,v1)
w <- fn.ab.2(length(qs),h=1000,r=1.5)</pre>
```

fn.ab.3

Vector of abundances by increasing root powers

# Description

Function to simulate haplotype abundances in the quasispecies by increasing root powers.

# Usage

```
fn.ab.3(n, h = 10000)
```

# **Arguments**

n Number of counts to compute.

h Highest abundance value.

# **Details**

The abundances, as counts, are computed according to the following equation, taking the integer part:

$$max(h^{(1/i)}, 1); i = 1..n$$

### Value

Numeric vector with n decreasing counts, where the first element equals h, and no element is lower than 1.

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### Author(s)

Josep Gregori and Mercedes Guerrero

### See Also

```
fn.ab.1, fn.ab.2, geom.series, GetRandomSeq, GenerateVars, Diverge
```

# **Examples**

```
# Simulate a quasispecies alignment.
m1 <- GetRandomSeq(50)
v1 <- GenerateVars(m1,50,2,c(10,1))
qs <- c(m1,v1)
w <- fn.ab.3(length(qs),h=1000)</pre>
```

FreqMat

Matrix of nucleotide or amino acid frequencies in alignment by position

# **Description**

Computes the nucleotide or amino acid frequency at each position in the alignment.

#### Usage

```
FreqMat(seqs,nr=NULL)
```

### **Arguments**

seqs DNAStringSet or AAStringSet with the aligned haplotype sequences.

An optional numeric vector with the haplotype counts.

#### Value

Matrix with the frequency of each nucleotide or amino acid in each position. A (4 x n) or (20 x n) matrix, where n is the alignment length.

#### Author(s)

Josep Gregori and Mercedes Guerrero

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Frequencies only in the alignment.
FreqMat(lst$hseqs)
# Also taking into account haplotype frequencies.
FreqMat(lst$hseqs,lst$nr)</pre>
```

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GenerateVars	Generate variants of a given haplotype	

# Description

Function to generate a set of variants for a given DNA sequence.

### Usage

```
GenerateVars(seq, nhpl, max.muts, p.muts)
```

### **Arguments**

seq A character string with a DNA sequence from which to generate the variants.

nhpl Number of haplotypes to generate.

max.muts Maximum number of mutations in each sequence.

p.muts Vector of length max.muts with the probability of each number of mutations,

some of which may be 0.

### **Details**

Given a DNA sequence, nhpl variant haplotypes are generated at random, with a maximum of max.muts substitutions each. The probability of the number of mutations in each haplotype generated is given by the vector p.muts. The positions of the mutations in each haplotype are independent and random.

# Value

A character vector with nhpl haplotype variants of the seq sequence.

# Author(s)

Josep Gregori and Mercedes Guerrero

#### See Also

```
GetRandomSeq, Diverge
```

```
set.seed(123)
m1 <- GetRandomSeq(50)
GenerateVars(m1,50,2,c(10,1))</pre>
```

GenotypeStandards\_A-H.fas

Genotype standards of hepatitis B virus

### **Description**

Fasta file with a set of well characterized sequences belonging to each HBV genotype. See the QSutils vignette: vignette("QSutils", package = "QSutils")

# **Format**

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

### See Also

DBrule

### **Examples**

```
lstRefs <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA") \\ RefSeqs <- lstRefs\$hseq
```

geom.series

Geometric series

# Description

Function to simulate haplotype abundances in the quasispecies by geometric series.

### Usage

```
geom.series(n,p=0.001)
```

### **Arguments**

n Number of frequencies to compute.

p Numeric parameter of the geometric function.

# **Details**

The abundances, as counts, are computed according to the following equation:

$$p(1-p)^{i-1}, i=1..n$$

The lower r, the faster the decrease in abundances.

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# Value

Numeric vector with n decreasing counts.

# Author(s)

Josep Gregori and Mercedes Guerrero

### See Also

```
fn.ab.1,fn.ab.2, fn.ab.3, GetRandomSeq, GenerateVars, Diverge
```

# **Examples**

```
# Simulate a quasispecies alignment.
m1 <- GetRandomSeq(50)
v1 <- GenerateVars(m1,50,2,c(10,1))
qs <- c(m1,v1)
w <- geom.series(100,0.8)</pre>
```

GetInfProfile

Information content profile of an alignment

# **Description**

GetInfProfile computes the information content at each position of an alignment.

# Usage

```
GetInfProfile(seqs,nr=NULL)
```

### **Arguments**

seqs DNAStringSet or AAStringSet with the haplotype alignment.

nr An optional numeric vector with the haplotype counts to take into account the

information content of each position in the alignment.

#### Value

Returns a numeric vector whose length is equal to the length of the alignment. Each value corresponds to the information content of each position in the alignment.

### Author(s)

Josep Gregori and Mercedes Guerrero

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### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

# **Examples**

```
# Load the alignment
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Compute the alignment's IC profile.
GetInfProfile(lst$hseqs)
# Also taking into account haplotype frequencies.
GetInfProfile(lst$hseqs,lst$nr)</pre>
```

GetQSData	Read the aligned sequences, filter at minimum abundance, and sort the sequences

# **Description**

Reads aligned amplicon sequences with abundance data, filters at a given minimum abundance, and sorts by mutations and abundance.

# Usage

```
GetQSData(flnm,min.pct=0.1,type="DNA")
```

# **Arguments**

flnm	Fasta file with haplotype sequences and their frequencies. The header of each haplotype in the fasta file is composed of an ID followed by a vertical bar "I" followed by the read counts, and eventually followed by another vertical bar and additional information (i.e. Hpl.2.0001 15874 25.2)
min.pct	Minimum abundance, in %, to filter the reads. Defaults to 0.1%.
type	Character string specifying the type of the sequences in the fasta file. This must be one of "DNA" or "AA". It is "DNA" by default.

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#### **Details**

The fasta file is loaded and the haplotype abundances, as counts, are taken from the header of each sequence. Haplotypes with abundances below min.pct % are filtered out. The haplotypes are then sorted: first, by decreasing order of the number of mutations with respect to the dominant haplotype, and second, by decreasing order of abundances. The haplotypes are then renamed according to the pattern Hpl.n.xxxx, where n represents the number of mutations, and xxxx the abundance order within the mutation number.

### Value

Returns a list with three elements.

bseqs DNAStringSet or AAStringSet with the haplotype sequences.

nr Vector of haplotype counts.

nm Vector of number of mutations of each haplotype with respect to the dominant

(most frequent) haplotype.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

# See Also

ReadAmplSeqs

```
lst<-GetQSData("QSutils/extdata/ToyData_10_50_1000.fna",min.pct=0.1,type="DNA")
lst</pre>
```

GetRandomSeq 21

GetRandomSeq

Generate a random sequence

# Description

Creates a random DNA sequence of a given length.

# Usage

```
GetRandomSeq(seq.len)
```

# Arguments

seq.len

The sequence length.

### Value

A character string representing a DNA sequence.

### Author(s)

Josep Gregori and Mercedes Guerrero

### See Also

GenerateVars, Diverge

# **Examples**

```
set.seed(123)
GetRandomSeq(50)
```

GiniSimpson

Functions to calculate the GiniSimpson index

# Description

GiniSimpson calculates the unbiased estimator, GiniSimpsonVar computes Gini-Simpson asymptotic variance, and GiniSimpsonMVUE calculates the minimum variance unbiased estimator of the Gini-Simpson index.

# Usage

```
GiniSimpson(w)
GiniSimpsonMVUE(w)
GiniSimpsonVar(w)
```

22 HCq

### **Arguments**

W

Vector of observed counts or frequencies.

#### Value

A value that corresponds to the Gini-Simpson diversity index.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

# Examples

```
# A vector of haplotype counts.
nr <- c(464, 62, 39, 27, 37, 16, 33, 54, 248, 20)
# Gini-Simpson index.
GiniSimpson(nr)
# Gini-Simpson variance.
GiniSimpsonVar(nr)
# MVUE Gini-Simpson index.
GiniSimpsonMVUE(nr)</pre>
```

**HCq** 

Set of functions to compute the Havrda-Charvat estimator

# **Description**

HCq computes the Havrda-Charvat estimator, and HCqVar computes the Havrda-Charvat asymptotic variance for a given exponent. By using HCqProfile, a Havrda-Charvat estimator is calculated for a predefined vector of exponents to obtain the full profile in the range, 0 to Inf.

### **Usage**

```
HCq(w, q)
HCqVar(w, q)
HCqProfile(w, q = NULL)
```

HCq 23

### Arguments

- w Vector of observed counts or frequencies.
- g Exponent. By default, a vector of values 1, 2, 3, 4 and Inf.

#### **Details**

In HCq only the first element in q is considered. HCqProfile is vectorized and considers all elements in q. When q is NULL: in this case, a default vector is taken to obtain the full profile in the range 0 to Inf.

#### Value

A value that corresponds to the Havrda-Charvat estimator when HCq or HCqVar is used. A data frame with the Havrda-Charvat estimator for each exponent when HCqProfile is used.

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

Pavoine, S. (2005). M?thodes statistiques pour la mesure de la biodiversit?. UMR CNRS 5558 Biometrie et Biologie Evolutive.

#### See Also

```
Hill, Renyi
```

```
# A vector of observed counts.
nr<-c(464, 62, 39, 27, 37, 16, 33, 54, 248, 20)
# Havrda-Charvat estimator for q=4.
HCq(nr,4)
# Havrda-Charvat estimator variance for q=4.
HCqVar(nr,4)
# Prolife of Havrda-Charvat estimator for 0:4 and Inf.
HCqProfile(nr,c(0:4,Inf))</pre>
```

24 Hill

```
# Full profile.
HCqProfile(nr)
```

Hill

Hill numbers

# **Description**

Functions to compute Hill numbers. Hill computes the Hill number of a single q value. HillProfile computes Hill numbers for all elements in vector q.

#### Usage

```
Hill(w, q)
HillProfile(w, q = NULL)
```

### **Arguments**

w Vector of observed counts or frequencies.

q Exponent.

# **Details**

In Hill, only the first element in q is considered. HillProfile is vectorized and considers all elements in q.When q is NULL: in this case, a default vector is taken to obtain the full profile in the range, 0 to Inf.

#### Value

A value or vector of values corresponding to the Hill number estimators of passed exponents.

#### Author(s)

Josep Gregori and Mercedes Guerrero

# References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

HCq, Renyi

IntersectStrandHpls 25

### **Examples**

```
# Vector of observed counts.
nr<-c(464, 62, 39, 27, 37, 16, 33, 54, 248, 20)
# Hill numbers of order 2.
Hill(nr,2)
# Set of most common values.
HillProfile(nr,q=c(0:4,Inf))
# Full Hill numbers profile.
HillProfile(nr)</pre>
```

IntersectStrandHpls

Forward and reverse strand haplotype intersections

# **Description**

Computes the intersection of forward and reverse strand haplotypes after a previous abundance filter that removes strand haplotypes below a given frequency threshold or unique to a single strand.

# Usage

```
IntersectStrandHpls(nrFW, hseqsFW, nrRV, hseqsRV, thr = 0.001)
```

# **Arguments**

nrFW	Numeric vector with forward strand haplotype counts.
hseqsFW	DNAStringSet object with the forward strand haplotypes.
nrRV	Numeric vector with forward reverse strand haplotypes.
hseqsRV	DNAStringSet object with the reverse strand haplotypes.
thr	Threshold to filter haplotypes at minimum abundance.

#### Value

List object with this elements:

hseqs DNAStringSet object with the forward and reverse strand intersected.

nr Numeric vector with the abundance of each haplotype.

pFW Vector of abundances of aligned forward strand.
PRV Vector of abundances of aligned reverse strand.

### Author(s)

Josep Gregori and Mercedes Guerrero

26 MutationFreq

### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

#### See Also

ReadAmplSeqs

#### **Examples**

```
# Load objects.
FW<- ReadAmplSeqs("QSutils/extdata/ToyData_FWReads.fna",type="DNA")
RV<- ReadAmplSeqs("QSutils/extdata/ToyData_RVReads.fna",type="DNA")
# Intersect the two objects, with a default threshold.
IntersectStrandHpls(FW$nr,FW$hseqs,RV$nr,RV$hseqs)</pre>
```

MutationFreq

Mutation frequency with respect to the dominant haplotype

### Description

MutationFreq computes the mutation frequency given a vector of counts, and the genetic distances of each haplotype to the dominant haplotype. MutationFreqVar returns the variance of the mutation frequency.

# Usage

```
MutationFreq(dst=NULL,nm=NULL,nr=NULL,len=1)
MutationFreqVar(nm,nr=NULL,len=1)
```

### **Arguments**

dst	A "dist" object or a symmetric matrix with pairwise distances.
nm	Vector of distances or differences with respect to the dominant haplotype including itself (eg, nm[1] is 0 if w[1]== $max(w)$ ).
nr	An optional numeric vector with the haplotype counts.
len	The alignment width when nm is the number of differences, otherwise 1. Defaults to 1

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#### Value

A value corresponding to the mutation frequency for MutationFreq or its variance for MutationFreqVar. When nr is NULL, the same weight is given to each haplotype and the computed value corresponds to the mutation frequency by entity.

#### Author(s)

Josep Gregori and Mercedes Guerrero

### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

#### See Also

```
DNA.dist, GetQSData, ReadAmplSeqs
```

# **Examples**

```
# Load alignment with abundances.
lst <- GetQSData("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")

# Mutation frequency.
dst <- DNA.dist(lst$seqs,model="raw")
MutationFreq(dst=dst,len=width(lst$seqs)[1])

# Mutation frequency with abundances.
MutationFreq(nm=lst$nm,nr=lst$nr,len=width(lst$seqs)[1])

# Variance of the mutation frequency.
MutationFreqVar(nm=lst$nm,nr=lst$nr,len=width(lst$seqs)[1])</pre>
```

MutsTbl

Table of mutation frequencies by position

### **Description**

Computes the table of mutation frequencies by position with respect to the alignment consensus.

### Usage

```
MutsTbl(hseqs,nr=NULL)
```

28 NucleotideDiversity

### **Arguments**

hseqs DNAStringSet or AAStringSet with the aligned haplotype sequences.

nr An optional numeric vector with the haplotype counts. When nr is NULL, the

same weight is given to each haplotype.

#### Value

Matrix of mutation counts by position. A (4 x n) or (20 x n) matrix, where n is the alignment length.

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

#### See Also

ReadAmplSeqs

### **Examples**

```
# Load the haplotypes alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Table of mutations in the alignment, regardless of haplotype abundance.
MutsTbl(lst$hseqs)
# Table of mutations taking into account abundance.
MutsTbl(lst$hseqs,lst$nr)</pre>
```

NucleotideDiversity Nucleotide diversity

#### **Description**

Computes the mean pairwise genetic distance between sequences in the alignment.

### Usage

NucleotideDiversity(dst,w=NULL)

NucleotideDiversity 29

# **Arguments**

dst	A "dist" object or a s	symmetrical matrix with	haplotype 1	pairwise distances (ie,

the output of DNA.dist).

w An optional numeric vector with the haplotype counts. When w is NULL, the

same weight is given to each haplotype, and nucleotide diversity is computed at

the entity level.

#### Value

A value that corresponds to the nucleotide diversity, either by entity or abundance, depending on argument w.

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

```
DNA.dist, ReadAmplSeqs
```

```
# Load haplotype alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Compute the DNA distance matrix.
dst <- DNA.dist(lst$hseqs,model="K80")
NucleotideDiversity(dst, lst$nr)
NucleotideDiversity(dst)</pre>
```

30 PolyDist

Po]	LvC	)is	st

Fraction of substitutions by polymorphic site

### **Description**

Computes the fraction of substitutions at each polymorphic site. The wild-type base is taken as the most abundant at each site, taking into account the weights, w.

### Usage

```
PolyDist(seqs, w=NULL)
```

### **Arguments**

seqs DNAStringSet or AAStringSet with the haplotype sequences.

w An optional numeric vector with the haplotype counts. When w is NULL, the

same weight is given to each haplotype.

#### Value

Vector of numbers corresponding to the fraction of substitutions at polymorphic sites. Note that the wild type also depends on w.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

ReadAmplSeqs

```
# Load haplotype alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
PolyDist(lst$hseqs)
PolyDist(lst$hseqs,lst$nr)</pre>
```

Rao 31

# **Description**

Set of functions to estimate Rao's functional entropy. Rao calculates the Rao entropy, RaoVar the variance of the Rao estimator, RaoPow the Rao entropy of order q, and RaoPowProfile the functional Rao entropy profile for the given set of exponents.

#### Usage

```
Rao(dst, w=NULL)
RaoVar(dst,w=NULL)
RaoPow(dst,q,w=NULL)
RaoPowProfile(dst,w=NULL,q=NULL)
```

# Arguments

dst	A "dist" object, output of the DNA.dist function.
W	An optional numeric vector with the haplotype counts. When w is NULL the same weight is given to each haplotype.
q	Exponent. A single value for Rao, RaoVar and RaoPow. A vector of values for RaoPowProfile. The default value for RaoPowProfile is a set of exponents to obtain a smooth profile.

### Value

A single value for Rao, RaoVar and RaoPow. A vector of values for RaoPowProfile corresponding to each exponent in vector q.

# Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

Pavoine, S. (2005). Méthodes statistiques pour la mesure de la biodiversité. UMR CNRS 5558 «Biométrie et Biologie Evolutive».

32 ReadAmplSeqs

### See Also

```
DNA.dist, ReadAmplSeqs
```

### **Examples**

```
# Load haplotype alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# DNA pairwise distances.
dst <- DNA.dist(lst$hseqs,model="N")

Rao(dst,lst$nr)
RaoVar(dst,lst$nr)
RaoPow(dst,2,lst$nr)
RaoPowProfile(dst,lst$nr,c(0:4,Inf))
RaoPowProfile(dst,lst$nr)</pre>
```

ReadAmplSeqs

Read a fasta file with haplotypes and frequencies

### Description

Loads an alignment of haplotypes and their frequencies from a fasta file.

### Usage

```
ReadAmplSeqs(flnm, type="DNA")
```

# Arguments

flnm File name of a fasta file with haplotype sequences and their frequencies. The

header of each haplotype in the fasta file is composed of an ID followed by a vertical bar "I" followed by the read count, and eventually followed by another

vertical bar and additional information (eg, Hpl.2.0001|15874|25.2).

type Character string specifying the types of sequences in the fasta file. This must be

either "DNA" or "AA". It is "DNA" by default.

### Value

Returns a list with two elements:

nr Vector of the haplotype counts.

hseqs DNAStringSet or AAStringSet with the haplotype DNA sequences or amino

acid sequences.

### Author(s)

Josep Gregori and Mercedes Guerrero

Renyi 33

### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

### See Also

GetQSData

#### **Examples**

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
lst</pre>
```

Renyi

Rényi profiles

### **Description**

Functions to compute the Rényi entropy given a vector of counts RenyiProfile computes the Rényi number for a set of exponents.

### Usage

```
Renyi(w, q)
RenyiProfile(w, q = NULL)
```

### Arguments

w Vector of observed counts or frequencies.

q Exponent. A single value for Renyi, a vector of values or NULL for RenyiProfile.

#### Value

A single value for Renyi. A data frame with exponents and entropies for RenyiProfile.

### Author(s)

Josep Gregori and Mercedes Guerrero

34 Report Variants

### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

Pavoine, S. (2005). Méthodes statistiques pour la mesure de la biodiversité. UMR CNRS 5558 «Biométrie et Biologie Evolutive».

### See Also

```
Hill, HCq
```

# **Examples**

```
# A vector of observed counts.
nr<-c(464, 62, 39, 27, 37, 16, 33, 54, 248, 20)
Renyi(nr,2)
RenyiProfile(nr,c(0:4,Inf))
RenyiProfile(nr)</pre>
```

ReportVariants

Report variants

# Description

eports the variants of a DNAStringSet or AAStringSet of haplotypes given a reference sequence.

# Usage

```
ReportVariants(hseqs,ref.seq,nr=NULL,start=1)
```

### **Arguments**

hseqs	DNAStringSet or AAstringSet object of the aligned haplotypes.
ref.seq	Character vector with the reference sequence of the alignment.
nr	Numeric vector with the abundances of each haplotype in hseqs. When nr is NULL, a vector of ones is taken as default.
start	Position of the first nucleotide in the alignment

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#### Value

A dataframe with 4 columns: the nucleotide in the reference sequence, the position, the variant nucleotide, and its abundance.

### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

### **Examples**

```
# Load objects.
lst<-ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Report the variants in these haplotypes,
# taking as a reference the most abundant haplotype.
ReportVariants(lst$hseqs[-1], ref.seq= as.character(lst$hseqs[1]),
lst$nr[-1], start = 1)</pre>
```

SegSites

Compute the number of segregating sites

### **Description**

Computes the number of segregating (polymorphic) sites in a given alignment. That is, the number of sites with more than a single nucleotide or amino acid in the alignment.

# Usage

```
SegSites(seqs)
```

### **Arguments**

seqs

DNAStringSet or AAStringSet with the haplotype sequences.

36 Shannon

### Value

A value corresponding to the number of polymorphic sites.

#### Author(s)

Josep Gregori and Mercedes Guerrero

### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

# See Also

ReadAmplSeqs

#### **Examples**

```
# Create the object.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
SegSites(lst$hseqs)</pre>
```

Shannon

Set of functions to compute Shannon entropy

# Description

Shannon computes the Shannon entropy. NormShannon returns the normalized Shannon entropy. ShannonVar computes the Shannon entropy asymptotic variance. NormShannonVar computes the normalized Shannon entropy asymptotic variance.

# Usage

```
Shannon(w)
ShannonVar(w)
NormShannon(w)
NormShannonVar(w)
```

# Arguments w

Vector of observed counts or frequencies.

SortByMutations 37

### Value

A single value with the result of the computations.

# Author(s)

Josep Gregori and Mercedes Guerrero

# References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

# **Examples**

```
# Create a vector of observed counts.
nr<-c(464, 62, 39, 27, 37, 16, 33, 54, 248, 20)
# Shannon entropy.
Shannon(nr)
# Shannon entropy variance.
ShannonVar(nr)
# Normalized Shannon entropy.
NormShannon(nr)
# Normalized Shannon entropy variance.
NormShannonVar(nr)</pre>
```

SortByMutations

Sort haplotypes by mutations and abundance

# **Description**

Sorts and renames haplotypes by the number of mutations with respect to the dominant haplotype, and by abundance.

# Usage

```
SortByMutations(bseqs, nr)
```

38 SortByMutations

### Arguments

bseqs DNAStringSet or AAStringSet object with the haplotype alignment.

nr Vector with the haplotype counts.

### **Details**

The haplotypes are pairwise-aligned to the dominant haplotype and then sorted: first, by decreasing order of the number of differences with respect to the dominant haplotype, and second, by decreasing order of abundance. As a result, haplotypes are renamed according to the pattern Hpl.n.xxxx, where n represents the number of differences, and xxxx the abundance order within the mutation number.

#### Value

Returns a list with three elements.

bseqs DNAStringSet or AAStringSet with the haplotype sequences.

nr Vector of the haplotype counts.

nm Vector of the number of differences of each haplotype with respect to the domi-

nant haplotype.

#### Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

ReadAmplSeqs

```
# Load haplotype alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
SortByMutations(lst$hseq,lst$nr)</pre>
```

SummaryMuts 39

SummaryMuts Distribution of nucleotides or amino acids in polymorphic sites
---

### **Description**

Computes the nucleotide or amino acid frequencies at all polymorphic sites in the alignment.

### Usage

```
SummaryMuts(seqs, w = NULL, off = 0)
```

# Arguments

5	eqs	DNAStringSet or AAStringSet with the haplotype sequences.	
٧	ı	An optional numeric vector with the haplotype counts. When w is NULL, a vector of ones is taken as default.	
c	off	Offset of first position in the alignment	

#### Value

Data frame with the polymorphic positions and nucleotide or amino acid frequencies.

### Author(s)

Josep Gregori and Mercedes Guerrero

### References

Gregori J, Esteban JI, Cubero M, Garcia-Cehic D, Perales C, Casillas R, Alvarez-Tejado M, Rodríguez-Frías F, Guardia J, Domingo E, Quer J. Ultra-deep pyrosequencing (UDPS) data treatment to study amplicon HCV minor variants. PLoS One. 2013 Dec 31;8(12):e83361. doi: 10.1371/journal.pone.0083361. eCollection 2013. PubMed PMID: 24391758; PubMed Central PMCID: PMC3877031.

Ramírez C, Gregori J, Buti M, Tabernero D, Camós S, Casillas R, Quer J, Esteban R, Homs M, Rodriguez-Frías F. A comparative study of ultra-deep pyrosequencing and cloning to quantitatively analyze the viral quasispecies using hepatitis B virus infection as a model. Antiviral Res. 2013 May;98(2):273-83. doi: 10.1016/j.antiviral.2013.03.007. Epub 2013 Mar 20. PubMed PMID: 23523552.

#### See Also

ReadAmplSeqs

40 TotalMutations

### **Examples**

```
# Load haplotype alignment with abundances.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
# Distribution of nucleotides at polymorphic sites.
SummaryMuts(lst$hseqs,lst$nr,off=0)</pre>
```

TotalMutations

Number of Mutations

# Description

TotalMutations computes the number of mutations in the alignment.

### Usage

TotalMutations(hseqs,w)

### **Arguments**

hseqs DNAStringSet or AAStringSet with the haplotype sequences.

An optional numeric vector with the haplotype counts used to compute the total

number of mutations in the population, that is, taking into account haplotype

abundances. When w is NULL, a vector of ones is taken as default.

### Value

A value corresponding to the number of mutations. Note that the wild-type is decided taking winto account.

# Author(s)

Josep Gregori and Mercedes Guerrero

#### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

### See Also

SegSites

Toy.GapsAndNs.fna 41

### **Examples**

```
# Create the object.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")

TotalMutations(lst$hseqs)
TotalMutations(lst$hseqs,lst$nr)</pre>
```

Toy.GapsAndNs.fna

Fasta file with raw reads with gaps and Ns

### **Description**

Fasta file of sequenced data with some missing information. This is toy data to illustrate some functions of the package QSutils.

#### **Format**

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

### See Also

Collapse, CorrectGapsAndNs and Recollapse

```
ToyData_10_50_1000.fna
```

Fasta file with 10 haplotypes, 50 basepairs in size.

### **Description**

Fasta file that contains the sequence of 10 haplotypes used as examples in the QSutils package.

#### **Format**

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

### **Examples**

```
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
lst</pre>
```

ToyData\_FWReads.fna

Fasta file with forward reads

### **Description**

Fasta file with forward strand reads. Toy data used to illustrate the intersections of forward and reverse haplotypes with the function IntersectStrandHpls.

#### Format

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

### See Also

ToyData\_RVReads.fna, IntersectStrandHpls

```
lstFW <- ReadAmplSeqs("QSutils/extdata/ToyData_FWReads.fna",type="DNA")
lstRV <- ReadAmplSeqs("QSutils/extdata/ToyData_RVReads.fna",type="DNA")
lstI <- IntersectStrandHpls(lstFW$nr,lstFW$hseqs,lstRV$nr,lstRV$hseqs)
lstI</pre>
```

ToyData\_RVReads.fna

Fasta file with reverse reads.

### **Description**

Fasta file with reverse strand reads. Toy data used to illustrate the intersections of forward and reverse haplotypes with the function IntersectStrandHpls.

### **Format**

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

#### See Also

```
ToyData_FWReads.fna, IntersectStrandHpls
```

### **Examples**

```
lstFW <- ReadAmplSeqs("QSutils/extdata/ToyData_FWReads.fna",type="DNA")
lstRV <- ReadAmplSeqs("QSutils/extdata/ToyData_RVReads.fna",type="DNA")
lstI <- IntersectStrandHpls(lstFW$nr,lstFW$hseqs,lstRV$nr,lstRV$hseqs)
lstI</pre>
```

UniqueMutations

Number of unique mutations

# **Description**

UniqueMutations computes the number of unique mutations in the alignment.

# Usage

UniqueMutations(hseqs)

# **Arguments**

hseqs

DNAStringSet or AAStringSet with the haplotype sequences.

#### Value

A value corresponding to the number of mutations.

### Author(s)

Josep Gregori and Mercedes Guerrero

### References

Gregori J, Perales C, Rodriguez-Frias F, Esteban JI, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016 Jun;493:227-37. doi: 10.1016/j.virol.2016.03.017. Epub 2016 Apr 6. Review. PubMed PMID: 27060566.

Gregori J, Salicrú M, Domingo E, Sanchez A, Esteban JI, Rodríguez-Frías F, Quer J. Inference with viral quasispecies diversity indices: clonal and NGS approaches. Bioinformatics. 2014 Apr 15;30(8):1104-1111. Epub 2014 Jan 2. PubMed PMID: 24389655.

#### See Also

TotalMutations

### **Examples**

```
# Create the object.
lst <- ReadAmplSeqs("QSutils/extdata/ToyData_10_50_1000.fna",type="DNA")
UniqueMutations(lst$hseqs)</pre>
```

Unknown-Genotype.fna Fasta file with reads of unknown genotype

### Description

Fasta file with hepatitis B virus sequences of unknown genotype. This is used to illustrate the genotyping of HBV sequences with the QSutils package.

# Format

Fasta file format. Each sequence starts with the symbol ">" followed by the sequence ID. Subsequent lines correspond to the nucleotide sequences or peptide sequences.

### See Also

DBrule

```
lst2Geno <- ReadAmplSeqs("QSutils/extdata/Unknown-Genotype.fna",type="DNA")
hseq <- lst2Geno$hseq[1]
hseq</pre>
```

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